Biocatalytic protein membranes fabricated by electrospinning

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Abstract

In this study, a protein-based catalytic membrane was produced by electrospinning. Membrane activity was characterised in terms of response current for various glucose concentrations. We focused on the preparation of a scaffold by converting a globular protein to other structural forms using catastrophic solvents. A scaffolding protein, bovine serum albumin, and an enzyme, glucose oxidase (GOD), were selected as a model natural carrier matrix and a biologically active agent, respectively. Beta-mercaptoethanol (β-ME) was used to convert the globular protein to an amyloid-like form. A structural stabilising agent, 2,2,2-trifluoroethanol (TFE), was used to maintain the final α-helical structure of the amyloid-like protein. The TFE:PBS (phosphate-buffered saline) ratio and various electrospinning parameters were analysed to minimise activity loss. Using this approach, we applied electrospinning to an active enzyme to obtain biocatalytic nanofibrous membranes. After optimising the protein electrospinning process, the activities of the protein nanofibrous membranes were monitored. GOD remained active in the new membrane structure. The highest enzyme activity was observed for the membranes prepared with a 1.5:1 (v:v) TFE:PBS solvent ratio. In that particular case, the immobilised enzyme created a current of 0.7 μA and the apparent activity was 2547 ± 132 U/m².

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1. Introduction

Electrospinning is a popular technique to produce nanofibrous structures using polymeric substances for a variety of applications [1–5]. The power of electrospinning mainly depends on the applicability of the technique to a number of different polymers with minor optimisation steps. Using the appropriate optimisation conditions, one can adjust the thickness, morphology, and length of a desired polymeric structure. Additionally, some studies have recently examined the limitation [7,8]. Additionally, some studies have recently examined direct electrospinning of proteins to form biological nanofibres [9]. Globular proteins, such as albumin [9–12], haemoglobin [5], and myoglobin [13], have been electrospun.

Our inspiration for protein electrospinning arose from the natural biological fibres secreted by diverse organisms. Biological protein fibres are generally secreted by specialised instruments, as in the case of the secretion of silk by Bombyx mori [14,15]. Silk is one of the oldest known protein fibres with functional properties. It has previously been reported that spider silk contains amyloid-like fibres when it is assembled in vitro [16]. Although biological fibres have many applications, the production of these natural fibres is not cost effective or efficient [17,18]. In this context, the utilisation of electrospinning to produce protein-based natural fibres is promising. Using this approach, different proteins with various functionalities can be electrospun for biotechnology applications. One of the best methods for fabricating a biofunctional scaffold is to load a bioactive agent to the scaffold. The performance of the bioactive agent is strongly dependent on supporting structures at the nanoscale. For example, nanofibres minimise diffusion limitations [9]. However, owing to their structural arrangement, the electrospinning of proteins requires an intensive optimisation strategy. As mentioned in the work of Dror and coworkers for the electrospinning of bovine serum albumin (BSA), disulphide bonds should be broken to obtain an electrospinnable solution using a catastrophic solvent. This process causes the conversion of a protein into an amyloid-like form. Among various alcohols, beta-mercaptoethanol (β-ME) is mainly used...
to open the tertiary structure of proteins and α-helical structures are stabilised by 2,2,2-trifluoroethanol (TFE) [10].

In this article, a BSA-derived amyloid-like structure was used as a natural polymer for the electrospinning process to enhance the coating and supporting properties during the entrapment of a bioactive agent. In this scenario, the glucose oxidase (GOD) enzyme was chosen as a model bioactive agent. The negative effects of catastrophic solvents on enzymatic activity were studied. The ratio of the solvent to stabilising agent was carefully adjusted, as were the electrospinning parameters to minimise activity loss. The preliminary results for the performance of electrospun biocatalytic membranes are reported based on apparent enzyme activity.

2. Experimental

2.1. Materials

Glucose oxidase (EC 1.1.3.4.) from Aspergillus niger (specific activity: 306 U/mg), TFE, β-ME, D-(+)-glucose monohydrate, and Congo red were purchased from Sigma Chemical Co. (St. Louis, MO, USA). All other reagents (NaCl, Na2HPO4, and NaH2PO4) used as buffering agents and BSA were supplied by Acros Organics (Morris Plains, NJ, USA). Double-distilled water was used throughout the experiments.

2.2. Preparation of solutions and electrospinning

The negative effect of catastrophic solvents on enzymatic activity was studied. BSA was dissolved in a mixture of TFE and phosphate-buffered saline (PBS) at a concentration of 12% (w/v) at room temperature. The tertiary structure of the protein was decomposed with 10 equiv. bond β-ME and stabilised by the addition of TFE at the following ratios: 1.5:1:0, 3.0:1:0, 4.5:1:0, and 9.0:1:0. The BSA-containing solvent was stirred continuously at 25 °C for 4 h. After obtaining a clear solution of protein, the powdered enzyme corresponding to 200 U/mL was added, stirred for 5 min, and the final solution was transferred to an injector to begin the electrospinning process immediately. Accordingly, the interaction period for the enzyme, GOD, and catastrophic solvents was limited to diminish activity prior to the electrospinning process. The preparation protocol for the solution is summarised in Fig. 1.

For the electrospinning setup, a direct current voltage supplier (MCH 303D2; Gamma High Voltage Research Inc., Ormond Beach, FL, USA) and a syringe pump (NE-1000; New Era Pump Systems Inc., Farmingdale, NY, USA) were used. The mixed solution was loaded into a 5-mL syringe and a syringe pump was located above the collector. The proportional flow rates (Q) were maintained between 0.20 and 0.60 mL/h (Fig. 1). Fibres were collected on glass slides and aluminium foil. The applied voltage and the distance between the tip and the collector varied between 9 and 20 kV and 10–20 cm, respectively. All experiments were performed at room temperature (about 25 °C). The samples were then stored for 24 h in a vacuum desiccator.

2.3. Characterisation

Fiber morphologies were investigated by optical microscopy (Nikon Eclipse LV100, Nikon Instruments Inc., Melville, NY, USA) and scanning electron microscopy (SEM, FEI-Quanta 200; Hillsboro, OR, USA). The chemical surfaces of membranes were characterised by FTIR-ATR (Fourier transform infrared spectroscopy attenuated total reflectance) (Perkin Elmer, Waltham, MA, USA). After production, electrospun membranes were stained with Congo red using the methods by Puchtler et al. [19] in order to verify the formation of an amyloid-like structure [19].

2.4. Measurement of enzyme activity

The reaction sequence to monitor glucose utilisation of flavin adenine dinucleotide (FAD) is given as follows:

\[
\beta-D-Glucose + \text{GOD(FAD)} \rightarrow \beta-D-Gluconolactone + \text{GOD(FADH}_2) \tag{1}
\]

\[
\text{GOD(FADH}_2) + O_2 \rightarrow \text{GOD(FAD)} + H_2O_2 \tag{2}
\]

\[
H_2O_2 \rightarrow O_2 + 2H^+ + 2e^- \tag{3}
\]

Apparent enzyme activity was measured by amperometric detection. Amperometric measurements were performed using the DropSens μStat 200 (Llanera, Spain). The DropSens electrode was polarised at 650 mV to decompose hydrogen peroxide (H2O2), which is formed during the reaction of glucose with oxygen in the presence of GOD (1, 2). The electrons that result from the decomposition of hydrogen peroxide are proportional to the glucose concentration (3). They were tracked amperometrically and the response current (I) was recorded in units of µA. The details of the amperometric measurements of enzymatic activity were described previously [20].

In the experimental setup, the electrospun membranes were placed in a 50-mL PBS solution and stirred for 10 min until equilibrium was reached. A stock glucose solution (200 mM) was added to the PBS solution to achieve a final concentration of 0.05–50 mM glucose, and the solution was stirred continuously. Enzyme activities corresponding to different substrate concentrations were investigated based on response currents. This procedure was repeated several times until deviations from linearity were observed for all membranes. The optimum solvent...
ratio was determined by the response current corresponding to highest apparent enzymatic activity.

Additionally, the effect of GOD content in electrospun membranes was investigated. Specifically, 200 U/mL, 100 U/mL, and 50 U/mL GOD were added to the solution. The membrane with the highest activity (Sample A) was electrospun for different time intervals (1–60 min) in order to observe the effects of accumulation period on the total activity of the membrane. Resulting current values with respect to glucose concentration for “Sample A” were converted to apparent enzymatic activity using a calibration curve. Furthermore, amperometric measurements were repeated after 24 h for membranes kept in PBS to examine the degradation of enzyme activity.

3. Results and discussion

3.1. Preparation and characterisation of protein-based biocatalytic membranes

Protein electrospinning requires an optimisation strategy. A basic problem with the utilisation of proteins for electrospinning is their structural stability and closed loop structures compared to synthetic polymers. In this study, we performed a tedious optimisation procedure to determine the correct conditions for protein electrospinning. A common alpha helix structural stabiliser, TFE, causes damage to the structure and function of proteins [21]. To determine the effect of TFE concentration on fibre morphology and enzyme activity, four solutions were prepared. In a preliminary analysis, enzymatic activity in solutions with different TFE:PBS ratios was detected and the relative enzymatic activities of the solutions were as follows: Sample A > Sample B > Sample C > Sample D.

Electrospinning parameters were optimised for the solution that had the lowest TFE:PBS ratio (i.e., Sample A), but the highest enzyme activity. First, the effect of applied voltage was examined. We assumed that protein-based membranes are more susceptible to higher voltages than synthetic polymers. Previous studies have examined the effect of higher voltages, but these results were not applicable to our optimisation study, which examined large-scale variation in voltage values [22–25]. In our study, electrospinning was carried out using 9–20 kV to evaluate the effect of voltage. High voltages (> 12 kV) caused ruptures because the fibres were thinner, and lower voltages (< 12 kV) resulted in thicker fibres. The threshold voltage value for the spinning process was 9 kV to obtain a considerable Taylor cone structure. SEM images of the fibres produced using these parameters are shown in Fig. 2A1 and Fig. 2A2. The nanofibres that were electrospun at 9 kV had an average diameter of 612 ± 64 nm, and those that were electrospun at 12 kV had an average diameter of 474 ± 23 nm. Keeping the voltage at 9 kV, despite the large fibre diameter, uniform fibres were observed. As a result, 9 kV was chosen as the working voltage. Secondly, the effect of flow rate on fibre morphology was studied. Electrospinning was performed for flow rates of 0.20–0.60 mL/h. As the flow rate increased, the solvent became less prone to evaporation and the diameters of the resultant fibres increased. When the flow rate was greater than 0.50 mL/h, more than one jet (unstable Taylor cone) was observed. The fibres that were electrospun with a flow rate of 0.50 mL/h had larger fibre diameters (1096 ± 124 nm); however, those that were electrospun with a flow rate of 0.35 mL/h had smaller fibre diameters (632 ± 23 nm) and were more uniform. When the flow rate was less than 0.35 mL/h, a stable Taylor cone could not form. Consequently, 0.35 mL/h was chosen as the working flow rate. Only the SEM images of fibres produced with flow rates of 0.35 and 0.50 mL/h are shown in Fig. 2B1 and Fig. 2B2. Finally, the effect of the tip-to-collector distance was examined. When the distance was less than 10 cm, the solvent did not evaporate and a fibrous network could not form. When the distance was greater than 20 cm, the jet did not reach the collector. The effects of distances between 10 and 20 cm were evaluated. SEM images of fibres produced with distances of 8 and 12 cm to the collector are shown in Fig. 2C1 and Fig. 2C2, respectively. The membrane that was electrospun with a distance of 8 cm had an average diameter of 742 ± 89 nm, and the membrane that was electrospun with a 12-cm distance had an average diameter of 323 ± 18 nm. The fibres shown in Fig. 2C1 are non-uniform and thick owing to the lack of solvent evaporation, which results from the small distance (8 cm). In contrast, the membranes shown in Fig. 2C2 are more uniform and thinner because the distance was increased.

Accordingly, the working parameters were optimised at 12 kV, 10 cm, and 0.35 mL/h for the solution with the lowest TFE concentration (Sample A). The other solutions (Samples B, C, and D) were further electrospun under these optimised conditions. The SEM images of all
samples are shown in Fig. 3A–D. The nanofibres of Sample A showed uniform sizes with the smallest diameters (227 ± 19 nm). However, nanofibres became less uniform as the TFE concentration increased (Samples B and C). The nanofibres of Sample B were uniform, but had thicker structures (552 ± 45 nm) compared to the fibres produced using Sample A. A drastic deformation of fibre structure was observed for membranes prepared with Sample C. Finally, the membrane prepared with Sample D, with the highest TFE concentration, had a gel-like structure because the solvent was not fully evaporated.

In order to demonstrate the amyloid-like formation of BSA in the electrospinning process, fibres stained with Congo red dye were observed under an optical microscope. As is clearly shown in Fig. 4, electrospun nanofibres sustained amyloid-like characteristics compared with nanofibres produced with the polycaprolactone polymer as a negative control. The amyloid-specific dye Congo red showed differences in amyloid membrane colour, while the polycaprolactone membrane did not show any difference after staining. Optical images of all protein fibres prepared using different parameters supported the amyloid-like form of the proteins (BSA and GOD) in the solutions [16].

Based on FTIR-ATR measurements, the amyloid-like structure of nanofibres was investigated. A broad peak between 1600 and 1700 cm\(^{-1}\) indicated the presence of an amide I band. In the case of GOD, the amide I band was the characteristic peak. The amide I band showed that the protein secondary structure included random coil and \(\alpha\)-helical conformations. As shown in Fig. 5, there was an obvious shift in the amide I band from 1646 cm\(^{-1}\) to 1660 cm\(^{-1}\) (from GOD to BSA–GOD). This may indicate the formation of a \(\beta\)-turn/antiparallel \(\beta\)-sheet conformation in the structure, i.e. the formation of an amyloid-like structure. There was also peak shift in the amide I band between BSA and BSA–GOD. These results showed that some GOD was converted to the amyloid-like form in the presence of catastrophic solvents [26].

3.2. Enzyme activity studies

3.2.1. Effect of TFE concentration on enzyme activity

The highest apparent enzyme activity was recorded for the fibres produced with the solution containing the lowest TFE concentration (Sample A). For Sample B and Sample C, the lower limit of enzyme
activity detection was 0.1 mM glucose; however, linearity was attained for 1.5 mM and 2 mM glucose, respectively. These conditions resulted in a limited detection interval. The lowest activity was recorded for the fibres produced with the solution containing the highest TFE concentration (Sample D), as expected. For Sample A, the linear range of glucose detection was 35 mM glucose; however, the lower limit of detection for glucose was 2 mM.

For the membranes prepared using Sample D, the enzyme activity was so low that no response was observed for glucose concentrations above 5 mM; however, the limit of detection for Sample D was as low as 0.05 mM glucose. The membranes prepared by Sample D can be used for the detection of low blood glucose levels owing to the low detection limit. However, the membranes prepared with Sample A can be used for normal and diabetic glucose detection, for which a larger detection interval is necessary.

3.2.2. Effect of enzyme concentration

The effect of enzyme concentration on enzyme activity was detected for the membranes prepared using Sample A, which had the highest activity in the previous analysis. The concentration of GOD did not affect the response current, substantially. These results can be explained by a diffusion-controlled mechanism of the glucose–GOD reaction in the electrospun nanofibre as well as the maximum enzymatic response upon severe destruction by TFE [27,28]. The effect of different GOD concentrations on enzyme activity is illustrated in Fig. 7.

The maximum currents were 0.72 μA for the membrane with 200 U/mL enzymatic activity, 0.65 μA for the membrane with 100 U/mL enzymatic activity, and as low as 0.30 μA for the membrane with 50 U/mL enzymatic activity. It is speculated that as the amount of enzyme decreases, the current also decreases due to the reduction in total enzymatic activity. Another explanation for the decrease in current could be the limited diffusion of the substrate.

Such enzymatic reactions have been examined extensively in our research for more than 20 years. For example, the reaction of glucose with GOD in various conditions was intensively examined in our biosensor studies. In a preliminary analysis, the diffusion of GOD in a biological matrix, porcine mucus, was examined [29]. In another study, we reported the diffusion-limited reaction kinetics of GOD and glucose for sandwich-type enzyme electrode applications [30]. Further studies were carried out using cross-linked matrices [31]. The diffusion of glucose over the enzyme layers formed in different conditions affects the reaction kinetics as well as the performance of the enzyme electrode [32–33].

![FTIR-ATR Spectra of BSA-GOD nanofibres, BSA fibres and GOD.](image)

Fig. 5. FTIR-ATR Spectra of BSA-GOD nanofibres, BSA fibres and GOD.

3.2.3. Effect of accumulation time

The effect of accumulation time was assessed using membranes prepared with Sample A and the results are shown in Fig. 8. It is quite clear that a longer period of fibre formation on the surface was associated with higher enzyme deposition on the surface. Accordingly, it is expected that as the accumulation time increases, the response or enzymatic activity will be higher. However, all response curves for membranes against glucose concentration showed a similar behaviour (Fig. 6). The membrane with the highest enzymatic activity was the one with an accumulation time of 1 min. The membrane with 60 min of accumulation time, which had the greatest enzyme content, showed slightly less activity than the 1-minute sample. The samples with accumulation times of 10, 20, and 40 min showed less activity than the 60-minute and 1-minute samples. These responses can only be explained by mass transfer limitations. In the experiments to obtain the response with respect to glucose concentration, each data point for every single enzymatic reaction was collected in not less than 10 min. However, these data were collected for free enzymes in calibration curve experiments within a few seconds, and not for more than 30 s. Therefore, it is quite clear that the system is controlled by the diffusion of the substrate, glucose, through the amyloid matrix. The movements of the substrate (glucose) to the enzyme and the product (hydrogen peroxide) to the electrode surface require more time than the reaction kinetics over the enzyme surface. This very slow diffusion–reaction–diffusion mechanism is directly related to the “effectiveness factor” and “Thiele modulus” of the system and will be investigated in our forthcoming research, similar to our previous studies [17]. Similarly, the response current values of the membrane prepared with the conditions used for “Sample A” were converted to apparent enzymatic activity by the calibration curve. The highest attainable response was observed for the membrane collected at 1 min, and the corresponding enzyme activity was 2547 ± 132 U/m².

4. Conclusion

In this study, a biocatalytic nanofibrous membrane was successfully prepared by electrospinning. First, the damage caused by TFE was minimised by making the protein solution electrospinnable with a 1.5:1.0 (v:v) TFE:PBS ratio. For this ratio, fibres showed more uniform structures with smaller diameters compared to other samples prepared.
with higher TFE:PBS ratios. The enzymatic activity for this membrane was also greater than that of the others, which proved that it was the most favourable membrane for biocatalytic applications. The effect of electrospinning parameters on membrane morphology was determined by changing the applied voltage, flow rate, and distance. Below 9 kV, a Taylor cone did not form and above 12 kV, ruptures were observed. For a flow rate of less than 0.35 mL/h, no Taylor cone formed and above 2.50 mL/h, the Taylor cone was unstable. If the distance between the tip and the collector was less than 10 cm, nanofibres did not form and above 20 cm, the jet was not able to reach the collector. Enzyme activity did not increase proportionally with accumulation time owing to mass transfer limitations. Similarly, the increase in enzyme activity was highly limited when the enzyme content in the membrane was doubled. This study provides a simple and efficient technique for entrapping an enzyme inside protein nanofibres; the method has several applications and can be used to develop biosensors in the future.

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